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Patentanmeldung Nr. Patent application No. Demande de brevet n°

99402236.6

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**Blatt 2 der Bescheinigung
Sheet 2 of the certificate
Page 2 de l'attestation**

Anmeldung Nr.: 99402236.6
Application no.:
Demande n°:

Anmeldetag: 10/09/99
Date of filing:
Date de dépôt:

Anmelder:
Applicant(s):
Demandeur(s):
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SWEDEN

Bezeichnung der Erfindung:

Title of the invention:

Titre de l'invention:

Variants of the gamma chain of AMPK, DNA sequences encoding the same, and uses thereof

In Anspruch genommene Priorität(en) / Priority(ies) claimed / Priorité(s) revendiquée(s)

Staat:
State:
Pays:

Tag:
Date:
Date:

Aktenzeichen:
File no.
Numéro de dépôt:

Internationale Patentklassifikation:
International Patent classification:
Classification internationale des brevets:

C12N15/54, C12N15/11, C12N9/12, C12Q1/68, A01K67/027, G01N33/68

Am Anmeldetag benannte Vertragsstaaten:
Contracting states designated at date of filing: AT/BE/CH/CY/DE/DK/ES/FI/FR/GB/GR/IE/IT/LI/LU/MC/NL/PT/SE/TR
Etats contractants désignés lors du dépôt:

Bemerkungen:
Remarks:
Remarques:

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VARIANTS OF THE GAMMA CHAIN OF AMPK, DNA SEQUENCES
ENCODING THE SAME, AND USES THEREOF.

The present invention relates to new variants
of the γ chain of AMP-activated protein kinase (AMPK), to
5 genes encoding said variants and to uses thereof.

AMPK has a key role in regulating the energy
metabolism in the eukaryotic cell (HARDIE et al., Annu.
Rev. Biochem., 67, 821-855, 1998; KEMP et al., TIBS, 24,
22-25, 1999). Mammalian AMPK is a heterotrimeric complex
10 comprising a catalytic α subunit and two non-catalytic β
and γ subunits that regulate the activity of the α
subunit. The yeast homologue (denoted SNF1) of this
enzyme complex is well characterised; it comprises a
catalytic chain (Snf1) corresponding to the mammalian α
15 subunit, and regulatory subunits: Sip1, Sip2 and Gal83
correspond to the mammalian β subunit, and Snf4
correspond to the mammalian γ subunit. Sequence data show
that AMPK homologues exist also in *Caenorhabditis elegans*
and *Drosophila*.

20 It has been observed that mutations in yeast
SNF1 and *SNF4* cause defects in the transcription of
glucose-repressed genes, sporulation, thermotolerance,
peroxisome biogenesis, and glycogen storage.

In the mammalian cells, AMPK has been proposed
25 to act as a "fuel gauge". It is activated by an increase
in the AMP:ATP ratio, resulting from cellular stresses
such as heat shock and depletion of glucose and ATP.
Activated AMPK turns on ATP-producing pathways (e.g.
fatty acid oxidisation) and inhibits ATP-consuming
30 pathways (e.g. fatty acid and cholesterol synthesis),
through phosphorylation of the enzymes acetyl-CoA
carboxylase and hydroxymethylglutaryl-CoA (HMG-CoA)
reductase. It has also been reported to inactivate in
vitro glycogen synthase, the key regulatory enzyme of
35 glycogen synthesis, by phosphorylation (HARDIE et al.,

1998, *supra*); however, whether glycogen synthase is a physiological target of AMPK *in vivo* remained unclear.

Several isoforms of the three different AMPK subunits are present in mammals. In humans, *PRKAA1* on human chromosome (HSA) 5p12 and *PRKAA2* on HSA1p31 respectively encode isoforms $\alpha 1$ and $\alpha 2$ of the α subunit, *PRKAB1* on HSA12q24.1 and *PRKAB2* (not yet mapped) respectively encode isoforms $\beta 1$ and $\beta 2$ of the β subunit, and *PRKAG1* on HSA12q13.1 and *PRKAG2* on HSA7q35-q36 respectively encode isoforms $\gamma 1$ and $\gamma 2$ of the γ subunit (OMIM database, <http://www.ncbi.nlm.nih.gov/omim/>, July 1999). HARDIE et al., [1998, *supra*] also mention the existence of a third isoform ($\gamma 3$) of the γ subunit of AMPK but do not provide any information about it. Analysis of the sequences of these γ subunits shows that they are essentially composed of four cystathione β synthase (CBS) domains whose function is unknown. No phenotypic effect resulting from a mutation in either of the AMPK subunits has yet been documented.

On the other hand, it has been observed that most Hampshire pigs have a high intramuscular glycogen concentration. In these pigs, glycogenolysis which occurs after slaughtering leads to an important decrease of the pH, resulting in acid meat having a reduced water-holding capacity and giving a reduced yield of cured cooked ham.

The locus (named *RN*) associated with high muscular content of glycogen was first identified by family segregation analysis of phenotypic data from Hampshire pigs (LE ROY et al., Genet. Res., 55, 33-40, 1990). A fully dominant allele, *RN*, correlated with high glycogen content occurs at a high frequency in most Hampshire populations while pigs from other breeds are assumed to be homozygous for the normal, recessive *rn*⁺ allele. Subsequent studies showed that *RN* carriers have a large increase (about 70%) of glycogen in skeletal muscle

but not in liver (MONIN et al., in 38th ICoMST, Clermont-Ferrand, FRANCE, 1992).

5 The large difference in glycogen content between RN^- and rn^+ pigs leads to marked differences in meat quality and technological yield (ENFÄLT et al., J. Anim. Sci., 75, 2924-2935, 1997). The RN^- allele is therefore of considerable economical significance in the pig industry and most breeding companies would like to reduce or eliminate this dominant mutation.

10 The RN phenotype can be determined by measuring the glycolytic potential in muscle biopsies from live animals, or after slaughter (MONIN et al., Meat Science, 13, 49-63, 1985). However, this method has severe limitations for application in practical breeding
15 programs. The accuracy of the test is not 100%: as there is some overlap in the phenotypic distribution of RN^- and rn^+ , the test is not able to distinguish RN^-/RN^- homozygotes and RN^-/rn^+ heterozygotes. Further, the sampling of muscle biopsies on live animals is invasive
20 and costly.

Thus, there is a strong need for the development of a simple diagnostic DNA test for the RN locus. Moreover, the dramatic phenotypic effect of the RN gene in pigs implies that this gene has an important role
25 in the regulation of carbohydrate metabolism in skeletal muscle in other vertebrates, in particular mammals.

Skeletal muscle and liver are the two major reservoirs of glycogen in mammals and the observation of an increased muscular glycogen while liver glycogen is
30 normal suggests that the RN^- phenotype maybe due to a mutation in a gene expressed in muscle but not in liver. The inventors have previously reported that the RN gene is located on pig chromosome 15 (MILAN et al., Mamm. Genome, 7, 47-51, 1996; MARIANI et al., Mamm. Genome, 7,
35 52-54, 1996; LOOFT et al., Genetics Selection Evolution, 28, 437-442, 1996). They have now discovered that the RN^-

allele is associated with a non-conservative mutation in a gene encoding a new muscle-specific isoform of the AMP-activated protein kinase (AMPK) γ chain.

5 The various aspects of the present invention are based upon the discovery and characterisation of this mutation and the identification and isolation of the mutant gene.

According to the invention it is shown that a mutation in a γ chain of AMPK results in an altered
10 regulation of carbohydrate metabolism, demonstrating that AMPK is an essential component of said metabolism. It is also provided a nucleic acid sequence encoding a muscle-specific isoform of the γ chain of AMPK. Thus it is provided means to regulate carbohydrate metabolism, more
15 specifically to detect and/or correct potential or actual dysfunctions of the regulation of carbohydrate metabolism, in particular in skeletal muscle.

The invention provides a polypeptide comprising an amino acid sequence having at least 70%
20 identity or at least 85% similarity, preferably 80% identity or at least 90% similarity, more preferably at least 90% identity or at least 95% similarity, and still more preferably at least 95% identity or at least 99% similarity, with the polypeptide SEQ ID NO: 2. The
25 invention also provides an isolated nucleic acid sequence encoding said polypeptide, as well as the complement of said nucleic acid sequence.

Said polypeptide represents a new muscle-specific isoform of the γ chain of AMPK, and will also be
30 hereinafter referred as Prkag3; the gene encoding said polypeptide will also be hereinafter referred as PRKAG3.

"Identity" of a sequence with a reference sequence refers to the percent of residues that are the same when the two sequences are aligned for maximum
35 correspondence between residues positions. A polypeptide having an amino acid sequence having at least X% identity

with a reference sequence is defined herein as a polypeptide whose sequence may include up to 100-X amino acid alterations per each 100 amino acids of the reference amino acid sequence. Amino acids alterations
5 include deletion, substitution or insertion of consecutive or scattered amino acid residues in the reference sequence.

"Similarity" of a sequence with a reference sequence refers to the percent of residues that are the
10 same or only differ by conservative amino acid substitutions when the two sequences are aligned for maximum correspondence between residues positions. A conservative amino acid substitution is defined as the substitution of an amino acid residue for another amino
15 acid residue with similar chemical properties (e.g. size, charge or polarity), which generally does not change the functional properties of the protein. A polypeptide having an amino acid sequence having at least X% similarity with a reference sequence is defined herein as
20 a polypeptide whose sequence may include up to (100-X) non-conservative amino acid alterations per each 100 amino acids of the reference amino acid sequence. Non-conservative amino acids alterations include deletion, insertion, or non-conservative substitution of
25 consecutive or scattered amino acid residues in the reference sequence.

For instance, searching the "GenBank nr" database using BLASTp (ALTSCHUL et al., Nucleic Acids Res., 25, 3389-3402, 1997) with default settings and the
30 whole sequence SEQ ID NO: 2 as a query, the higher percents of identity or similarity with SEQ ID NO: 2 were found for:

- γ 1 subunit of human AMPK: 65% identity or 82% similarity (score: 399);
- 35 - γ 1 subunit of rat AMPK: 65% identity or 82% similarity (score: 399);

- γ 1 subunit of murine AMPK: 64% identity or 80% similarity (score: 390);

- γ subunit of Drosophila AMPK: 53% identity or 75% similarity (score: 332);

5 - Yeast Snf4: 33% identity or 56% similarity (score: 173).

Polypeptides of the invention include for instance any polypeptide (whether natural, synthetic, semi-synthetic, or recombinant) from any vertebrate
10 species, more specifically from birds, such as poultry, or mammals, including bovine, ovine, porcine, murine, equine, and human, and comprising, or consisting of, the amino acid sequence of either:

- 15 - a functional Prkag3; or
- a functionally altered mutant of Prkag3.

"Functional" refers to a protein having a normal biological activity. Such a protein may comprise silent mutations inducing no substantial change in its activity, and having no noticeable phenotypic effects.
20 Non limitative examples of functional Prkag3 are:

- a porcine Prkag3 represented in the enclosed sequence listing under SEQ ID NO: 2;
25 - a human Prkag3 represented in the enclosed sequence listing under SEQ ID NO: 4;
- a V40I variant of Prkag3 resulting from the substitution of a valine residue in position 40 by an isoleucine residue.

A "functionally altered mutant" of a protein
30 comprises one or several mutations inducing a substantial change in its activity. Such mutations include in particular deletions, insertions, or non-conservative substitutions of amino acid residues in a domain essential for the biological activity of said protein.
35 They may result for instance in a partial or total loss of activity, or conversely in an increase of activity, or

in an impairment of the response to regulatory effectors. A non-limitative example of a functionally altered mutant of Prkag3 is the R41Q variant resulting from the non-conservative substitution of an arginine residue by a glutamine residue in position 41. This non-conservative substitution occurs inside a portion of the first CBS domain that is highly conserved between Prkag3 and the previously known isoforms of the γ subunit of AMPK.

Residue numbers for Prkag3 refer to the amino acid numbering of SEQ ID NO: 2 or SEQ ID NO: 4. Alignment of human and porcine Prkag3 sequences with previously known $\gamma 1$ and $\gamma 2$ isoforms is shown in Figure 3.

The invention also provides mutants of Prkag3 which may for instance be obtained by deletion of part of a Prkag3 polypeptide. Said mutants are generally functionally altered. They may have an identity with the overall Prkag3 sequence lower than 70%. However, the identity of the non-deleted sequences of said mutants, when aligned with the corresponding Prkag3 sequences should remain higher than 70%. Said mutants may for instance result from the expression of nucleic acid sequences obtained by deletion or insertion of a nucleic acid segment, or by a punctual mutation introducing a nonsense codon, in a nucleic acid sequence encoding a functional Prkag3.

The invention also provides a functionally altered mutant of a γ subunit of AMPK, wherein said mutant comprises at least one mutation responsible for said functional alteration located within the first CBS domain, and preferably within the region thereof aligned with the region spanning from residue 30 to residue 50 of a Prkag3 isoform. Said mutation may result from the insertion, deletion, and/or non-conservative substitution of one amino-acid or of several amino-acids, adjacent or not. More preferably the mutation is located within the region spanning from residue 35 to residue 45 of a Prkag3

isoform, or within the region of a $\gamma 1$ or $\gamma 2$ isoform, aligned therewith, for instance within the region spanning from residue 65 to residue 75 of the $\gamma 1$ isoform. According to a particular embodiment, said mutation is a non-conservative substitution, preferably a R→Q substitution. Advantageously, the mutation is located at residue 41 of the Prkag3 isoform, or at a corresponding residue of a $\gamma 1$ or $\gamma 2$ isoform, for instance at residue 70 of the $\gamma 1$ isoform.

The invention also provides a heterotrimeric AMPK wherein the γ subunit consists of a polypeptide of the invention.

The invention also provides isolated nucleic acid sequences encoding any of the above-defined functional or functionally altered Prkag3 or functionally altered mutants of a γ subunit of AMPK, and nucleic acid sequences complementary of any one of these nucleic acid sequences.

This includes particularly any isolated nucleic acid having the sequence of any of the naturally occurring alleles of a PRKAG3 gene, as well as any isolated nucleic acid having the sequence of an artificial mutant of a PRKAG3 gene, provided that said nucleic acid does not consist of the EST GENBANK AA178898.

This also includes any isolated nucleic acid having the sequence of a natural or artificial mutant of a PRKAG1 or a PRKAG2 gene, wherein said mutant encodes a functionally altered $\gamma 1$ or $\gamma 2$ subunit of AMPK as defined above.

Nucleic acid sequences of the invention may be obtained by the well-known methods of recombinant DNA technology and/or of chemical DNA synthesis. These methods also allow to introduce the desired mutations in a naturally occurring DNA sequence.

Examples of nucleic acid sequences encoding naturally occurring alleles of a *PRKAG3* gene are SEQ ID NO: 1, which encodes a naturally occurring allele of the porcine gene and SEQ ID NO: 3, which encodes a naturally occurring allele of the human gene. They may be used to generate probes allowing the isolation of *PRKAG3* from other species or of other allelic forms of *PRKAG3* from a same species, by screening a library of genomic DNA or of cDNA.

The invention also includes genomic DNA sequences from any vertebrate species, more specifically from birds, such as poultry, or mammals, including in particular bovine, ovine, porcine, murine, equine, and human, comprising at least a portion of a nucleic acid sequence encoding a polypeptide of the invention, preferably a portion of a *PRKAG3* gene, and up to 500 kb, preferably up to 100 kb of a 3' and/or of a 5' adjacent genomic sequence.

Such genomic DNA sequences may be obtained by methods known in the art, for instance by extension of a nucleic acid sequence encoding a polypeptide of the invention, employing a method such as restriction-site PCR (SARKAR et al., PCR Methods Applic., 2, 318-322, 1993), inverse PCR (TRIGLIA et al., Nucleic Acids Res., 16, 8186, 1988) using divergent primers based on a *Prkag3* coding region, capture PCR (LAGERSTROM et al., PCR Methods Applic., 1, 111-119, 1991), or the like.

The invention also includes specific fragments of a nucleic acid sequence encoding a polypeptide of the invention, or of a genomic DNA sequence of the invention as well as nucleic acid fragments specifically hybridising therewith. Preferably these fragments are at least 15bp long, more preferably at least 20bp long.

"Specific fragments" refers to nucleotidic sequences that are found only in the nucleic acids sequences encoding a polypeptide of the invention, and

are not found in nucleic acids sequences encoding related polypeptides of the prior art. This excludes the nucleic acid fragments that consist of a sequence shared with one of the known *PRKAG1* or *PRKAG2* genes.

5 "Specifically hybridising fragments" refers to nucleic acid fragments which can hybridise, under stringent conditions, only with nucleic acid sequences encoding a polypeptide of the invention, without hybridising with nucleic acid sequences encoding related
10 polypeptides of the prior art. This excludes the nucleic acid fragments that consist of the complement of a sequence shared with one of the known *PRKAG1* or *PRKAG2* genes.

Nucleic acid fragments that consist of the EST
15 GENBANK AA178898 or the complement thereof are also excluded.

Said specific or specifically hybridising nucleic acid fragments may for example be used as primers or probes or for detecting and/or amplifying a nucleic
20 acid sequence encoding a polypeptide of the invention. The invention encompasses set of primers comprising at least one primer consisting of a specific or specifically hybridising nucleic acid fragment as defined above.

The invention also provides recombinant
25 vectors comprising a nucleic acid sequence encoding a polypeptide of the invention. Vectors of the invention are preferably expression vectors, wherein a sequence encoding a polypeptide of the invention is placed under control of appropriate transcriptional and translational
30 control elements. These vectors may be obtained and introduced in a host cell by the well-known recombinant DNA and genetic engineering techniques.

The invention also comprises a prokaryotic or
eukaryotic host cell transformed by a vector of the
35 invention, preferably an expression vector.

A polypeptide of the invention may be obtained by culturing the host cell containing an expression vector comprising a nucleic acid sequence encoding said polypeptide, under conditions suitable for the expression
5 of the polypeptide, and recovering the polypeptide from the host cell culture.

A heterotrimeric AMPK wherein the γ subunit consists of a polypeptide of the invention may be obtained by expressing, together or separately, a nucleic
10 acid sequence encoding a polypeptide of the invention, a nucleic acid sequence encoding an α subunit, and a nucleic acid sequence encoding a β subunit, and reconstituting the heterotrimer.

The polypeptides thus obtained, or immunogenic
15 fragments thereof may be used to prepare antibodies, employing methods well known in the art. Antibodies directed against the whole Prkag3 polypeptide and able to recognise any variant thereof may thus be obtained. Antibodies directed against a specific epitope of a
20 particular variant (functional or not) of Prkag3 or antibodies directed against a specific epitope of a functionally altered mutant having a mutation in the first CBS domain of a γ subunit of AMPK, and able to recognise said variant or functionally altered mutant may
25 also be obtained.

As shown herein, mutations in a γ subunit of AMPK, and particularly mutations in the first CBS domain of a γ subunit of AMPK are likely to cause disorders in the energy metabolism (e.g. diabetes, obesity) in
30 vertebrates, including humans. Further, mutations in the first CBS domain or other parts of the PRKAG3 gene are likely to cause disorders in the muscular metabolism leading to diseases such as myopathy, diabetes and cardiovascular diseases.

35 The present invention provides means for detecting and correcting said disorders.

More specifically, the present invention is directed to methods that utilise the nucleic acid sequences and/or polypeptidic sequences of the invention for the diagnostic evaluation, genetic testing and prognosis of a metabolic disorder.

For example, the invention provides methods for diagnosing of metabolic disorders, more specifically carbohydrate metabolism disorders, and preferably disorders correlated with an altered, in particular an excessive, glycogen accumulation in the cells, resulting from a mutation in a gene encoding a γ subunit of AMPK, wherein said methods comprise detecting and/or measuring the expression of a functionally altered *PRKAG3* gene, or of a functionally altered mutant of a γ subunit of AMPK having a mutation within the first CBS domain in a nucleic acid sample obtained from a vertebrate, or detecting a mutation in the *PRKAG3* gene or in a sequence encoding the first CBS domain of a γ subunit of AMPK in the genome of a vertebrate suspected of having such a disorder.

According to a preferred embodiment of the invention, the disorder is correlated with an altered, in particular an excessive, glycogen accumulation in the muscular cells and results from the expression of a functionally altered *PRKAG3* gene.

The expression of a functionally altered *Prkag3*, or of a functionally altered mutant of a γ subunit of AMPK having a mutation within the first CBS domain may be detected or measured using either polyclonal or monoclonal antibodies specific for the functionally altered polypeptides of the invention, as defined above. Appropriate methods are known in the art. They include for instance enzyme-linked immunosorbent assay (ELISA), radioimmunoassay (RIA), and fluorescence activated cell sorting (FACS).

The nucleotide sequences of the invention may be used for detecting mutations in the *PRKAG3* gene or in a sequence encoding the first CBS domain of a γ subunit of AMPK, by detection of differences in gene sequences or in adjacent sequences between normal, carrier, or affected individuals.

The invention provides a process for detecting a mutation in the *PRKAG3* gene or in a sequence encoding the first CBS domain of a γ subunit of AMPK wherein said process comprises:

- obtaining a nucleic acid sample from a vertebrate;
- checking the presence in said nucleic acid sample of a nucleic acid sequence encoding a mutant *Prkag3*, or a mutant of a γ subunit of AMPK having a mutation within the first CBS domain, as defined above.

According to a preferred embodiment of the invention there is provided a method for detecting a nucleic acid sequence comprising a mutation in the *PRKAG3* gene or in a sequence encoding the first CBS domain of a γ subunit of AMPK wherein said process comprises:

- obtaining a nucleic acid sample from a vertebrate;
- contacting said nucleic acid sample with a nucleic acid probe obtained from a nucleic acid of the invention and spanning said mutation, under conditions of specific hybridisation between said probe and the mutant sequence to be detected;
- detecting the hybridisation complex.

Preferably, the process of the invention further comprises, prior to hybridisation, PCR amplification from the nucleic acid sample, of a sequence comprising at least the portion of the *PRKAG3* sequence or of the sequence encoding the first CBS domain of the γ subunit of AMPK wherein the mutation is to be detected.

Methods allowing the specific hybridisation of a probe only with a perfectly matching complementary sequence, and useful for the detection of punctual

mutations are known in the art. They include for instance Allele Specific PCR (GIBBS, Nucleic Acid Res., 17, 2427-2448, 1989), Allele Specific Oligonucleotide Screening (SAIKI et al., Nature, 324, 163-166, 1986), and the like.

5 A mutation in the *PRKAG3* gene may also be detected through detection of polymorphic markers closely linked to said mutation.

The invention also provides means for identifying said polymorphic markers, and more specifically polymorphic markers comprised within a genomic DNA sequence comprising at least a portion of a *PRKAG3* gene, and up to 500 kb, preferably 300 kb, more preferably up to 100 kb of a 3' and/or of a 5' adjacent sequence.

15 Said polymorphic markers may be obtained for instance, by screening a genomic DNA library from a vertebrate with a probe specific for the *PRKAG3* gene, in order to select clones comprising said nucleic acid sequence and flanking chromosomal sequences, and identifying a polymorphic marker in said flanking chromosomal sequences. The allele(s) of a polymorphic marker associated with a given mutant allele of the *PRKAG3* gene may also easily be identified by use of a genomic DNA library from an individual wherein the presence of said mutant allele has previously been detected by hybridisation with a nucleic acid probe of the invention.

Polymorphic markers include for instance, single nucleotide polymorphisms (SNP), microsatellites, insertion/deletion polymorphism and restriction fragment length polymorphism (RFLP). These polymorphic markers may be identified by comparison of sequences flanking the *PRKAG3* gene obtained from several individuals. Microsatellites may also be identified by hybridisation with a nucleic acid probe specific of known microsatellite motifs.

Once a polymorphic marker has been identified, a DNA segment spanning the polymorphic locus may be sequenced and a set of primers allowing amplification of said DNA segment may be designed.

5 The invention also encompasses said DNA primers.

 Detection of a mutation in the *PRKAG3* gene may be performed by obtaining a sample of genomic DNA from a vertebrate, amplifying a segment of said DNA spanning a
10 polymorphic marker by polymerase chain reaction using a set of primers of the invention, and detecting in said amplified DNA the presence of an allele of said polymorphic marker associated with said mutation.

 By way of example, polymorphic markers which
15 may be obtained according to the invention, and DNA primers allowing the detection of polymorphic markers closely linked to the *RN* allele of porcine *PRKAG3* gene are listed in Table 1 hereinafter.

 According to a preferred embodiment of the
20 invention, the vertebrate is a mammal, preferably a farm animal and more preferably a porcine, and the mutation to be detected produces a functionally altered *Prkag3*. The detection of said mutation allows to predict whether said mammal or the progeny thereof is likely to have a high
25 intramuscular glycogen concentration. An example of such a mutation produces a functionally altered *Prkag3* having a R41Q substitution.

 The present invention also includes kits for the practice of the methods of the invention. The kits
30 comprise any container which contains at least one specific fragment of a nucleic acid sequence of the invention, or at least one nucleic acid fragment able to specifically hybridise with a nucleic acid sequence of the invention. Said nucleic acid fragment may be
35 labelled. The kits may also comprise a set of primers of the invention. They may be used in conjunction with

commercially available amplification kits. They may also include positive or negative control reactions or markers, molecular weight size markers for gel electrophoresis, and the like.

5 Other kits of the invention may include antibodies of the invention, optionally labelled, as well as the appropriate reagents for detecting an antigen-antibody reaction. They may also include positive or negative control reactions or markers.

10 The invention further provides means for modulating the expression of vertebrate genes encoding a γ subunit of AMPK, and more specifically of the *PRKAG3* gene and/or the synthesis or activity of the products of said genes.

15 A purified AMPK heterotrimer comprising wild-type or mutant *Prkag3* subunit, or a functionally altered mutant γ subunit having a mutation in the first CBS domain, may be used for screening *in vitro* compounds able to modulate AMPK activity, or to restore altered AMPK activity. This may be done, for instance, by:

- measuring the binding of the compound to said heterotrimer, using for example high-throughput screening methods; or,
 - measuring changes in AMPK kinase activity,
- 25 using for example high-throughput screening methods.

High throughput screening methods are disclosed, for instance, in "High throughput screening: The Discovery of Bioactive Substances", J.P. DEVLIN (Ed), MARCEL DEKKER Inc., New York (1997).

30 Nucleic acids of the invention may be used for therapeutic purposes. For instance, complementary molecules or fragments thereof (antisense oligonucleotides) may be used to modulate AMPK activity, more specifically in muscular tissue.

Also, a nucleic acid sequence encoding a functional Prkag3 may be used for restoring a normal AMPK function.

Transformed cells or animal tissues expressing
5 a wild-type or mutant Prkag3, or a functionally altered mutant of a γ subunit of AMPK as defined above, or expressing an AMPK comprising said mutant Prkag3, or said functionally altered mutant of a γ subunit of AMPK, may be used as *in vitro* model for elucidating the mechanism of
10 AMPK activity or for screening compounds able to modulate the expression of AMPK.

The screening may be performed by adding the compound to be tested to the culture medium of said cells or said tissues, and measuring alterations in energy
15 metabolism in said cells or said tissues using methods such as measurements of glucose concentrations (levels), glucose uptake, or changes of the ATP/AMP ratio, glycogen or lipid/protein content.

The invention provides animals transformed
20 with a nucleic acid sequence of the invention.

In one embodiment, said animals are transgenic animals having at least a transgene comprising a nucleic acid of the invention.

In another embodiment, said animals are
25 knockout animals. "Knockout animals" refers to animals whose native or endogenous *PRKAG3* alleles have been inactivated and which produce no functional Prkag3 of their own.

In light of the disclosure of the invention of
30 DNA sequences encoding a wild-type or mutant Prkag3, or a functionally altered mutant of a γ subunit of AMPK, transgenic animals as well as knockout animals may be produced in accordance with techniques known in the art, for instance by means of *in vivo* homologous
35 recombination.

Suitable methods for the preparation of transgenic or knock-out animals are for instance disclosed in: *Manipulating the Mouse Embryo*, 2nd Ed., by HOGAN et al., Cold Spring Harbor Laboratory Press, 1994; 5 *Transgenic Animal Technology*, edited by C. PINKERT, Academic Press Inc., 1994; *Gene Targeting: A Practical Approach*, edited by A.L. JOYNER, Oxford University Press, 1995; *Strategies in Transgenic Animal Science*, edited y G.M. MONASTERSKY and J.M. ROBL, ASM Press, 1995; *Mouse* 10 *Genetics: Concepts and Applications*, by Lee M. SILVER, Oxford University Press, 1995.

These animals may be used as models for metabolic diseases and disorders, more specifically for diseases and disorders of glycogen metabolism in muscle. 15 For instance they may be used for screening test molecules. Transgenic animals may thus be used for screening compounds able to modulate AMPK activity. Knockout animals of the invention may be used, in particular, for screening compounds able to modulate 20 energy metabolism, more specifically carbohydrate metabolism, in the absence of functional Prkag3.

The screening may be performed by administering the compound to be tested to the animal, and measuring alterations in energy metabolism in said 25 animal using methods such as glucose tolerance tests, measurements of insulin levels in blood, changes of the ATP/AMP ratio, glycogen or lipid/protein content in tissues and cells.

Transgenic or knock-out farm animals with 30 modified meat characteristics or modified energy metabolism may also be obtained.

The present invention will be further illustrated by the additional description which follows, which refers to examples of obtention and use of nucleic 35 acids of the invention. It should be understood however that these examples are given only by way of illustration

of the invention and do not constitute in any way a limitation thereof.

EXAMPLE 1: ISOLATING THE PRKAG3 GENE

We have screened a porcine Bacterial
5 Artificial Chromosome (BAC) library (ROGEL-GAILLARD et
al., Cytogenet and Cell Genet, 851, 273-278, 1999) and
constructed a contig of overlapping BAC clones across the
region of pig chromosome 15 harbouring the RN gene. These
BAC clones were in turn used to develop new genetic
10 markers in the form of single nucleotide polymorphisms
(SNPs) or microsatellites (MS) as described in Table 1
below.

Table 1

	Name of marker	BAC clone	Primer sequences	Size of PCR product (bp)	Marker type ^a	Alleles ^b
1	H3	115B9, 156E6, 361B4, 90A9	F: 5'-GGAAATTTCAAGTCAGCCAAAC-3' (SEQ ID NO: 5) R: 5'-CTTCAAAAGACCGTGCTACT-3' (SEQ ID NO: 6)	114 - 138	MS	114, 126, 128, 132*, 134*, 136, 138
2	MS982H1	982H11	F: 5'-CTGGGAACCTCTATATGCTG-3' (SEQ ID NO: 7) R: 5'-TAGGGAATACAAATCACAG-3' (SEQ ID NO: 8)	114 - 157	MS	114, 140, 142*, 144, 146, 150, 158
3	MS479L3	479L3, 297D7, 852B5, 153B5	F: 5'-CTCCAGCTCACAGGATGACA-3' (SEQ ID NO: 9) R: 5'-GTTTCTGCAGCTTAGCATCTATTCC-3' (SEQ ID NO: 10)	150 - 164	MS	150*, 160, 162, 164
4	MS997M3	997F12	F: 5'-GAAGTATCCTGGGCTTCTGA-3' (SEQ ID NO: 11) R: 5'-GTTTCTCCAGGTTCCAGACATCCAC-3' (SEQ ID NO: 12)	138 - 160	MS	138, 144, 152, 154, 160*
5	MS482H6	482E7	F: 5'-GCTTCTGTCTGCCCTACTT-3' (SEQ ID NO: 13) R: 5'-GTTTCTAAGTTCTACTGTAGACACC-3' (SEQ ID NO: 14)	78 - 90	MS	78, 80, 88*, 90
6	MS337H2	808G10, 947E5, 337G11	F: 5'-CCAAGCTGTGGTGGCTGAAT-3' (SEQ ID NO: 15) R: 5'-CAGCACAGCAGTGCCACCTA-3' (SEQ ID NO: 16)	145 - 165	MS	145, 149, 155, 161*, 165*
7	MS127B1	127G6, 134C9	F: 5'-CAAACCTCTTAGGCGTGT-3' (SEQ ID NO: 17) R: 5'-GTTTCTGGAATCCATATGCCATGG-3' (SEQ ID NO: 18)	94 - 108	MS	94, 100, 108*, 114
8	CMKAR2	128A3, 337G11, 808G10, 947E7, 1110H12	F: 5'-AGGGTGGATGGTAGGCTTCA-3' (SEQ ID NO: 19) R: 5'-GTCTCGCTCCTGAAGGAAGT-3' (SEQ ID NO: 20)	208	SNP	112A*, 112T; 158A*, 158G 176A*, 176G
9	127G63	127G6, 134C9, 170D7, 1030A5, 1088F2	F: 5'-AGTCACGTGGCCATGCTATC-3' (SEQ ID NO: 21) R: 5'-CTCAACTGGATTGAGTCAGT-3' (SEQ ID NO: 22)	409	SNP	234A*, 234C
10	VIL1		F: 5'-TTGGCGCAACTGTTATTTCT-3' (SEQ ID NO: 23) R: 5'-AGGCAAAAGGAAGACACAG-3' (SEQ ID NO: 24)	270	SNP	90T, 90G, 120A, 120G, 166C, 166T
11	NRAMP1	315F7, 530A6, 651C12, 1088F2, 1095H3	F: 5'-AGCCGTGGGCATCGTTGG-3' (SEQ ID NO: 25) R: 5'-AGAAGGAGACAGACAGGGCGA-3' (SEQ ID NO: 26)	1300	RFLP (Styl)	1: 100+1200 bp 2: 100+200+1000 bp

^aMS=microsatellite; SNP=single nucleotide polymorphism.^bMicrosatellite alleles are designated according to the length of the amplified fragment while SNPs are denoted according to the polymorphic nucleotide. Alleles associated with the *RN* allele are marked with an asterisk.

The new markers were used together with some previously described markers to construct a high-resolution linkage map. Standard linkage analysis using pedigree data comprising about 1,000 informative meioses for segregation at the RN locus made it possible to exclude RN from the region proximal to MS479L3 and distal to microsatellite Sw936. Linkage Disequilibrium (LD) analysis was done with the same markers and a random sample of 68 breeding boars from the Swedish Hampshire population, scored for the RN phenotype by measuring glycogen content in muscle. The results of LD analysis using the DISMULT program (TERWILLIGER, Am. J. Hum. Genet., 56, 777-787, 1995) are shown in Figure 1. They reveal a sharp LD peak around the markers MS127B1 and SNP127G63. These markers appeared to show complete linkage disequilibrium with the RN allele, i.e. RN was associated with a single allele at these two loci. The most simple interpretation of this finding is that the RN mutation arose on a chromosome carrying these alleles and that the two markers are so closely linked to the RN locus that the recombination frequency is close to 0%. The two markers are both present on the overlapping BAC clones 127G6 and 134C9 suggesting that the RN gene may reside on the same clone or one of the neighbouring clones.

A shot-gun library of the BAC clone 127G6 was constructed and more than 1,000 sequence reads were collected giving about 500,000 base pair random DNA sequence from the clone. The data were analysed and sequence contigs constructed with the PHRED, PHRAP and CONSED software package (University of Washington Genome Center, <http://bozeman.mbt.washington.edu>). The sequence data were masked for repeats using the REPEATMASKER software (<http://ftp.genome.washington.edu/cgi-bin/RepeatMasker>) and BLAST searches were carried out using the NCBI web site (<http://www.ncbi.nlm.nih.gov>).

Three convincing matches to coding sequences were obtained. Two of these were against human cDNA sequences/genes, KIAA0173 described as being similar to pig tubulin-tyrosine ligase and located on HSA2q (UniGene cluster Hs.169910, <http://www.ncbi.nlm.nih.gov/UniGene/>) and CYP27A1 located on HSA2q33-ter (UniGene cluster Hs.82568). The results strongly suggested that the pig coding sequences are orthologous to these human genes as it is well established that the RN region is homologous to HSA2q33-36 (ROBIC et al., Mamm. Genome, 10, 565-568, 1999). However, none of these sequences appeared as plausible candidate genes for RN. The third coding sequence identified in BAC 127G6 showed highly significant sequence similarity to various AMP-activated protein kinase γ sequences including the yeast *SNF4* sequence. The cDNA sequence of this gene was determined by RT-PCR and RACE analysis using muscle mRNA from an *rn⁺/rn⁺* homozygote. This sequence is shown in Figure 2 and in the enclosed sequence listing under SEQ ID NO: 1.

Legend of Figure 2:

5' UTR: 5' untranslated region

3' UTR: 3' untranslated region

CDS: coding sequence

***: stop codon

'-': identity to master sequence

'.' : alignment gap

The frame of translation was determined on the basis of homology to other members in the protein family and assuming that the first methionine codon in frame is the start codon. The deduced polypeptidic sequence is shown in the enclosed sequence listing under SEQ ID NO: 2. Figure 3 shows an amino acid alignment constructed with the CLUSTAL W program (THOMPSON et al., Nucleic Acids Research, 22, 4673-4680, 1994) with representative AMPK γ sequences in the nucleotide databases.

Legend of Figure 3:

Sequences used:

- HumG1: Genbank U42412
 RatG1: Genbank X95578
 5 MusG1: Genbank AF036535
 HumG2: coding sequences extracted from genomic sequence
 Genbank AC006966
 PigG3: this study
 HumG3: this study
 10 Dros: Genbank AF094764 (only the region showing homology
 to AMPKG/SNF4 is included)
 SNF4 (yeast): Genbank M30470

Abbreviations:

- *: stop codon
 15 '-': identity to master sequence
 '.': alignment gap
 blank: no information
 ?: poor sequence data

- 20 Table 2 below shows the amino acid (above
 diagonal) and nucleotide sequence (below diagonal)
 identities (in %) among mammalian, *Drosophila* and yeast
 AMPKG/SNF4 sequences.

TABLE 2

	PigG3	HumG3	HumG1	RatG1	MusG1	HumG2	Dros	SNF4
PigG3	-	97.0	64.2	64.2	63.9	62.6	53.2	34.0
HumG3	90.7	-	63.6	63.6	63.6	62.6	53.5	34.4
HumG1	64.2	64.5	-	96.7	96.3	75.6	60.9	33.5
RatG1	65.8	65.8	88.0	-	97.4	75.3	61.1	33.5
MusG1	65.3	64.8	87.2	92.8	-	74.6	61.7	33.5
HumG2	61.6	61.6	68.1	67.8	65.9	-	63.1	34.5
Dros	58.4	58.4	59.0	59.3	59.0	60.0	-	36.2
SNF4	44.0	44.2	45.4	44.6	45.3	45.7	44.8	-

- 25 Figure 4 shows a Neighbor-Joining phylogenetic
 tree constructed with the PAUP software (SWOFFORD,
*Phylogenetic analysis using parsimony (and other
 methods)*, Sinauer Associates, Inc. Publishers,
 Sunderland, Massachusetts, 1998) using yeast SNF4 as
 outgroup; support for branch orders obtained in bootstrap

analysis with 1,000 replicates are indicated, scales of tree is indicated at the bottom. The result showed that the pig gene located in the RN region is distinct from mammalian *PRKAG1* and *PRKAG2* isoforms and most likely
5 orthologous to a human gene represented by the human EST sequence AA178898 (GenBank) derived from a muscle cDNA library. This gene is herein denoted *PRKAG3* since it is the third isoform of a mammalian AMP-activated protein kinase γ characterised so far.

10 The cDNA sequence of this gene was determined by RT-PCR and 5'RACE analysis using human skeletal muscle cDNA (Clontech, Palo Alto, CA) and the deduced protein sequence showed 97% identity to the porcine sequence (Figure 2; Table 2). The cDNA sequence of human *PRKAG3* is
15 also shown in the enclosed sequence listing under SEQ ID NO: 3; the deduced polypeptidic sequence is shown in the enclosed sequence listing under SEQ ID NO: 4.

Using the high resolution human TNG radiation hybrid
panel (<http://shgc-www.stanford.edu/RH/TNGindex.html>), we mapped the human
20 homologs of *PRKAG3*, *CYP27A1* and *KIAA0173*, all present in the porcine BAC127G6. The three genes are also very closely linked in the human genome. *PRKAG3* was mapped at a distance of 33 cR_{50.000} from *KIAA0173* and 52 cR_{50.000} from
25 *CYP27A1*, with lod score support of 6.8 and 4.5, respectively.

The established role of AMPK in regulating energy metabolism, including glycogen storage, and its location in the region showing maximum linkage disequilibrium made *PRKAG3* a very strong candidate gene
30 for RN. This was further strengthened by hybridisation analysis of a human multiple tissue northern blots (CLONTECH, Palo Alto, CA) using human *PRKAG1* (IMAGE clone 0362755 corresponding to GenBank entry AA018675), human
35 *PRKAG2* (IMAGE clone 0322735 corresponding to GenBank

entry W15439) and a porcine *PRKAG3* probe. The results are shown in Figure 5.

Legend of Figure 5:

H: Heart, B: Brain, Pl: Placenta, L: Lung,
5 Li: Liver, M: Skeletal muscle, K: Kidney, Pa: Pancreas,
S: Spleen, Th: Thymus, P: Prostate, T: Testis, O: Ovary,
I: Small intestine, C: Colon (mucosal lining),
PBL: Peripheral Blood Leukocyte.

While the *PRKAG1* and *PRKAG2* probes showed a
10 broad tissue distribution of expression, *PRKAG3* showed a
distinct muscle-specific expression. This result is also
supported by the human EST database where multiple ESTs
representing *PRKAG1* and *PRKAG2* have been identified in
various cDNA libraries whereas a single EST (GenBank
15 entry AA178898) representing *PRKAG3* has been obtained
from a muscle cDNA library. The muscle-specific
expression of *PRKAG3* and the lack of expression in liver
are entirely consistent with the phenotypic effect of *RN⁻*,
namely that glycogen content is altered in muscle but
20 normal in liver (ESTRADE et al., Comp. Biochem. Physiol.
104B, 321-326, 1993).

The entire coding sequence and parts of the
5'UTR and 3'UTR were determined from *rn⁺/rn⁺* and *RN⁻/RN⁻*
homozygotes by RT-PCR analysis. A comparison revealed a
25 total of seven nucleotide differences between the
sequence from *rn⁺* and *RN⁻* animals, as shown in Table 3
below. Two of these were missense mutations, three were
silent substitutions and two were single nucleotide
substitutions in the presumed 5'UTR region. The single
30 exon carrying the two missense mutations and two silent
substitutions was also PCR amplified using genomic DNA
from additional *rn⁺* and *RN⁻* pigs from different breeds.
This revealed four different *PRKAG3* haplotypes for codons
34-41 and only the R41Q missense substitution was
35 exclusively associated with *RN⁻*. The nonconservative
substitution R41Q occurs in the most conserved region

among isotypic forms of the AMPK γ chain and arginine at this residue (number 70 in Prkag1) is conserved among different isoforms of mammalian AMPK γ sequences as well as in the corresponding *Drosophila* sequence (Figure 3). A simple diagnostic DNA test for the R41Q mutation was designed based on the oligonucleotide ligation assay (OLA; LANDEGREN et al., Science, 241, 1077-1080, 1988). Screening a large number of RN⁻ and rn⁺ animals from the Hampshire breed as well as large number of rn⁺ animals from other breeds showed that the 41Q allele was present in all RN⁻ animals but not found in any rn⁺ animals, as shown in Table 4 below. The absence of the 41Q allele from other breeds is consistent with the assumption that the RN⁻ allele originated in the Hampshire breed; the allele has not yet been found in purebred animals from other breeds. In conclusion, the results provide convincing evidence that PRKAG3 is identical to the RN gene and that the R41Q substitution most likely is the causative mutation.

20

Table 3. Comparison of the *PRKAG3* sequences associated with the *rn*⁺ and *rn*⁻ alleles in different pig populations^a

Associated Allele	RN allele	5'UTR nt83 nt152	Codon							Population ^b
			34	35	40	41	213			
1	<i>rn</i> ⁻	C T	GCC Ala	CTG Leu	GTC Val	CAA Gln	TCT Ser	H		
2	<i>rn</i> ⁺	A C	--T -	T-- -	--- -	-G- Arg	--C -	H, L, LW, M, WB		
3	<i>rn</i> ⁺	N.D. N.D.	--T -	T-- -	A-- Ile	-G- Arg	N.D. N.D.	H, LW, WB		
4	<i>rn</i> ⁺	N.D. N.D.	--T -	--- -	--- -	-G- Arg	N.D. N.D.	L, LW, WB		

^anucleotide and codon numbers are defined in Figure 2^bH=Hampshire, L=Landrace, LW=Large White, M=Meishan, WB=Wild Boar

N.D.=not determined, "--" indicates identity to the top sequence.

TABLE 4.

RN phenotype	Genotype at nucleotide 122			Total
	A/A	G/A	G/G	
RN ⁻ , Hampshire ^a	40	87	0	127
RN ⁻ , Hampshire ^{a,b}	0	13	0	13
rn ⁺ , Hampshire ^a	0	0	60	60
rn ⁺ , other breeds ^c	0	0	488	488

^arepresent both French and Swedish Hampshire populations

^bheterozygosity RN/rn⁺ deduced using pedigree information

^cbreeds: Angler Saddleback, n=31; Blond Mangalitza, n=2; Bunte Bentheimer, n=16; Duroc, n=160; Göttinger Minipig, n=4; Landrace, n=83; Large White, n=72; Meishan, n=8; Piétrain, n=75; Red Mangalitza, n=5; Rotbunte Husumer, n=15; Schwalbenbauch Mangalitza, n=7; Schwäbisch Hällische, n=2; European Wild Boar, n=5; Japanese Wild Boar, n=3.

Without being bound to any particular mechanism, it may be hypothesised that the AMPK heterotrimer including PRKAG3 is involved in the regulation of glucose transport into skeletal muscle. It has recently been reported that AMPK activation induced by the AMP analogue AICAR or by muscle contraction leads to an increased glucose uptake in skeletal muscle (BERGERON et al., Am. J. Physiol., 276, E938-944, 1999; HAYASHI et al., Diabetes, 47, 1369-1373, 1998). If this is the function of the AMPK heterotrimer including PRKAG3, R41Q may be a gain-of-function mutation causing a constitutively active holoenzyme, for instance due to the loss of an inactivating allosteric site. An increased uptake of glucose to skeletal muscle is expected to lead to an increase in muscle glycogen content as observed in RN⁻ animals. It has been shown that overexpression of glucose transporter 4 (GLUT4) in transgenic mice leads to increased uptake of glucose and increased glycogen storage (TREADWAY et al., J. Biol. Chem., 269, 29956-29961, 1994). This type of gain-of-function model is consistent with the dominance of RN⁻ as the presence of a single unregulated copy would have a large effect on AMPK enzyme activity.

An alternative hypothesis on the functional significance of the R41Q substitution associated with the *RN* allele may also be proposed. Based on the established roles of the yeast SNF1 enzyme in utilisation of glycogen and of mammalian AMPK for inhibiting energy-consuming pathways and stimulating energy-producing pathways, activated AMPK is expected to inhibit glycogen synthesis and stimulate glycogen degradation. If this is the functional role of the isoform(s) containing the *PRKAG3* product, the R41Q substitution would be a loss-of-function mutation or a dominant-negative mutation locking the AMPK heterotrimer in an inactive state. In these cases the phenotypic effect should be explained by haplo-insufficiency, since *RN* appears fully dominant.

EXAMPLE 2: DETECTION OF THE R41Q SUBSTITUTION IN PIG *PRKAG3*

A part of *PRKAG3* including codon 41 was amplified in 10 µl reactions containing 100 ng genomic DNA, 0.2 mM dNTPs, 1.5 mM MgCl₂, 4.0 pmol of both forward (AMPKG3F3: 5'-GGAGCAAATGTGCAGACAAG-3') and reverse (AMPKG3R2: 5'-CCCACGAAGCTCTGCTTCTT-3') primer, 10% DMSO, 1 U of Taq DNA polymerase and reaction buffer (ADVANCED BIOTECH, London, UK). The cycling conditions included an initial incubation at 94°C for 5 min followed by 3 cycles at 94°C (1 min), 57°C (1 min) and 72°C (1 min), and 35 cycles of 94°C (20 sec), 55°C (30 sec) and 72°C (30 sec). Allele discrimination at nucleotide position 122 was done using the oligonucleotide ligation assay (OLA, LANDEGREN et al., Science, 241, 1077-1080, 1988). The OLA method was carried out as a gel-based assay. Each 10 µl OLA reaction contained 0.5 pmol of each probe SNPRN-A (5'-Hex-TGGCCAACGGCGTCCA-3'), SNPRN-G (5'-ROX-GGCCAACGGCGTCCG-3') and SNPRN-Common (5'-phosphate-AGCGGCACCTTTGTGAAAAAAAAA-3'), 1.5 U of thermostable AMPLIGASE and reaction buffer (EPICENTRE TECHNOLOGIES, Madison, WI) and 0.5 µl of the AMPKG3F3/AMPKG3R2 PCR product. After an initial

incubation at 95°C for 5 min, the following thermocycling profile was repeated 10 times: denaturation at 94°C (30 sec), and probe annealing and ligation at 55°C (90 sec). After OLA cycling, 1 µl of product was heat denatured at 94°C (3 min), cooled on ice, and loaded onto 6% polyacrylamide denaturing gel for electrophoresis on an ABI377 DNA sequencer (PERKIN ELMER, Foster City, USA). The resulting fragment lengths and peak fluorescence were analysed using GENESCAN software (PERKIN ELMER, Foster City, USA).

The OLA-based method for the R41Q mutation was used to determine the genotype of DNA samples collected from 68 Swedish Hampshire animals phenotyped as either RN⁻ or rn⁺ based on their glycolytic potential (GP) value. Figure 6 illustrates typical OLA results from the three possible genotypes. All RN⁻ animals were scored as homozygous A/A (n=28) or heterozygous A/G (n=36) at nucleotide position 122 whereas the rn⁺ animals were homozygous G/G (n=4) at this position.

EXAMPLE 3: PREDICTING THE PRESENCE OF THE RN⁻ ALLELE USING A CLOSELY LINKED MICROSATELLITE, MS127B1

A microsatellite 127B1 (MS127B1) was cloned from BAC 127G7 containing pig PRKAG3. The BAC clone was digested with *Sau3AI* and the restriction fragments subcloned into the *BamHI* site of pUC18. The resulting library was probed with a (CA)₁₅ oligonucleotide probe labelled with [γ -³²P]-dATP. Strongly hybridising clones were sequenced and primers for PCR amplification of microsatellite loci were designed. Ten µl PCR reactions were performed containing 100 ng genomic DNA, 0.2 mM dNTPs, 1.5 mM MgCl₂, 4.0 pmol of both forward (MS127B1F: 5'-Fluorescein-CAAACCTCTTCTAGGCGTGT-3') and reverse (MS127B1R: 5'-GTTTCTGGAACCTTCCATATGCCATGG-3') primers, and 1 U of Taq DNA polymerase and reaction buffer (ADVANCED BIOTECH, London, UK). The cycling conditions included an initial incubation at 94°C for 5 min followed by 3 cycles at 94°C

(1 min), 57°C (1 min) and 72°C (1 min), and 35 cycles of 94°C (20 sec), 55°C (30 sec) and 72°C (30 sec). The PCR products (0.3 µl) were separated using 4% polyacrylamide denaturing gel electrophoresis on an ABI377 DNA sequencer (PERKIN ELMER, Foster City, USA). The resulting fragment lengths were analysed using the GENESCAN and GENOTYPER software (PERKIN ELMER, Foster City, USA).

The method was used to determine the genotype of DNA samples collected from 87 Swedish Hampshire animals phenotyped as either RN⁻ or rn⁺ based on their glycolytic potential (GP) value. Allele 108 (bp) showed a complete association to the RN⁻ allele in this material as all RN⁻ (RN⁻/RN⁻ or RN⁻/rn⁺) animals were homozygous or heterozygous for this allele while no rn⁺ (rn⁺/rn⁺) animals carried this allele, as shown in Table 5 below.

Table 5

Animals	n	Genotype				
		94/94	94/108	94/114	100/108	108/108
RN ⁻	80	0	37	0	2	41
rn ⁺	7	3	0	4	0	0

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CLAIMS

1) A gamma subunit of a vertebrate AMP-activated kinase (AMPK), wherein said gamma subunit is a polypeptide having at least 70% identity with the polypeptide SEQ ID NO: 2.

2) A polypeptide of claim 1, wherein said polypeptide has at least 95% identity with the polypeptide SEQ ID NO:2.

3) A polypeptide of any of claims 1 or 2, wherein said polypeptide has the sequence SEQ ID NO: 2 or SEQ ID NO:4 or a sequence resulting from a V40I substitution in SEQ ID NO: 2.

4) A polypeptide which is a functionally altered mutant of a gamma subunit of a vertebrate AMP-activated kinase, wherein said polypeptide has at least a mutation located within the first CBS domain of said gamma subunit.

5) A polypeptide of claim 4, wherein the mutation is located within the region of the first CBS domain aligned with the region of a polypeptide of SEQ ID NO: 2 spanning from residue 30 to residue 50.

6) A polypeptide of claim 5, wherein the mutation is a R→Q substitution.

7) A polypeptide of claim 6, having a sequence resulting from a R41Q substitution in SEQ ID NO: 2.

8) A polypeptide which is a mutant of a gamma subunit of a vertebrate AMP-activated kinase, wherein said polypeptide results from a deletion of a part of a polypeptide of claim 1.

9) A nucleic acid sequence encoding a polypeptide of any of claims 1 to 8, or the complement thereof, provided that said nucleic acid sequence does not consist of the EST GENBANK AA178898.

10) A nucleic acid sequence of claim 9, having the sequence SEQ ID NO: 1 or SEQ ID NO:3, or the complement thereof.

11) A nucleic acid sequence comprising at least a portion of a nucleic acid sequence encoding a polypeptide of any of claims 1 to 8, and up to 500 kb of a 3' and/or of a 5' adjacent genomic DNA sequence, or the complement thereof.

12) A nucleic acid fragment selected among:

- a specific fragment of a nucleic acid sequence encoding a polypeptide of any of claims 1 to 8, or of a nucleic acid sequence of claim 11;
- a nucleic acid fragment which specifically hybridises under stringent conditions with a nucleic acid sequence encoding a polypeptide of any of claims 1 to 8, or of a nucleic acid sequence of claim 11;

provided that said nucleic acid fragment does not consist of the EST GENBANK AA178898.

13) A set of primers for amplifying a nucleic acid sequence of any of claims 9 to 11 or a portion thereof, comprising at least a primer consisting of a nucleic acid fragment of claim 12.

14) A recombinant vector comprising a nucleic acid sequence encoding a polypeptide of any of claims 1 to 8.

15) An host cell transformed by a nucleic acid sequence encoding a polypeptide of any of claims 1 to 8.

16) A transgenic animal transformed by a nucleic acid sequence encoding a polypeptide of any of claims 1 to 8.

17) A knockout animal, wherein the gene encoding a polypeptide of claim 1 is inactive.

18) A heterotrimeric AMPK wherein the γ subunit consists of a polypeptide of any of claims 1 to 8.

19) A method of detecting a metabolic disorder resulting from a mutation in a gene encoding a γ subunit of AMPK, wherein said process comprises:

- obtaining a nucleic acid sample from a vertebrate;

- checking the presence in said nucleic acid of a nucleic acid sequence encoding a polypeptide of any
5 of claims 1 to 8, wherein said polypeptide is functionally altered.

20) A method of claim 19 wherein the disorder is correlated with an altered glycogen accumulation in the muscular cells and results from the expression of a
10 functionally altered allele of a polypeptide of claim 1.

21) A method of any of claims 19 or 20 wherein the presence of the nucleic acid sequence encoding said mutant polypeptide is checked by contacting said nucleic acid sample with a nucleic acid probe obtained from a
15 nucleic acid of claim 12 and spanning said mutation, under conditions of specific hybridisation between said probe and the mutant sequence to be detected, and detecting the hybridisation complex.

22) A method for obtaining a pair of primers
20 allowing to detect a genetic polymorphic marker linked to a nucleic acid sequence encoding a polypeptide of claim 1, wherein said process comprises:

- screening a genomic DNA library from a vertebrate with a probe specific for a nucleic acid
25 sequence encoding a polypeptide of claim 1, in order to select clones comprising said nucleic acid sequence and flanking chromosomal sequences;

- identifying a polymorphic locus in said flanking chromosomal sequences, and sequencing a DNA
30 segment comprising said polymorphic locus ;

- designing primer pairs flanking said polymorphic locus.

23) A method of claim 22 wherein the selected clones comprise at least a portion of a nucleic acid
35 sequence encoding a polypeptide of claim 1, and up to 500 kb of a 3' and/or of a 5' adjacent sequence.

24) A method of any of claims 19 to 23 wherein the vertebrate is a mammal.

25) A method of claim 24 wherein said mammal is a pig.

5 26) A pair of primers obtainable by the process of any of claims 21 to 25.

27) A process for detecting a dysfunction of carbohydrate metabolism resulting from the expression of a functionally altered allele of a polypeptide of claim 1
10 in a vertebrate, wherein said process comprises:

- obtaining a sample of genomic DNA from said vertebrate;

- contacting said DNA with a pair of primers of claim 26 under conditions allowing PCR amplification;

15 - analysing the PCR product to detect if an allele of a polymorphic marker linked to a nucleic acid sequence encoding a functionally altered allele of a polypeptide of claim 1 is present.

28) A process of claim 27, wherein said
20 functionally altered polypeptide results from a R41Q substitution.

29) A process of any of claims 27 or 28, wherein said vertebrate is a mammal.

30) A process of claim 29 wherein said mammal
25 is a pig.

31) A process of claim 30 wherein the pair of primers is selected among:

- a pair of primers consisting of SEQ ID NO: 5 and SEQ ID NO: 6;

30 - a pair of primers consisting of SEQ ID NO: 7 and SEQ ID NO: 8;

- a pair of primers consisting of SEQ ID NO: 9 and SEQ ID NO: 10;

35 - a pair of primers consisting of SEQ ID NO: 11 and SEQ ID NO: 12;

- a pair of primers consisting of
SEQ ID NO: 13 and SEQ ID NO: 14;
- a pair of primers consisting of
SEQ ID NO: 15 and SEQ ID NO: 16;
- 5 - a pair of primers consisting of
SEQ ID NO: 17 and SEQ ID NO: 18;
- a pair of primers consisting of
SEQ ID NO: 19 and SEQ ID NO: 20;
- 10 - a pair of primers consisting of
SEQ ID NO: 21 and SEQ ID NO: 22;
- a pair of primers consisting of
SEQ ID NO: 23 and SEQ ID NO: 24;
- a pair of primers consisting of
SEQ ID NO: 25 and SEQ ID NO: 26.
- 15 32) Use of a transformed cell of claim 15 to
screen compounds able to modulate AMPK activity.
- 33) Use of a transgenic animal of claim 16 to
screen compounds able to modulate AMPK activity.
- 34) Use of a knockout animal of claim 17 to
20 screen compounds able to modulate energy metabolism in
the absence of a functional polypeptide of claim 1.
- 35) Use of an heterotrimeric AMPK of claim 18
to screen compounds able to modulate AMPK activity.

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ABSTRACT

The invention concerns variants of the gamma chain of vertebrate AMP-activated kinase (AMPK), as well as nucleic acid sequences encoding said variants and use thereof for the diagnosis or treatment of dysfunction of energy metabolisms.

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LOOFT, CHRISTIAN
KALM, ERNST

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Figure 1. Linkage disequilibrium in the *RN* region

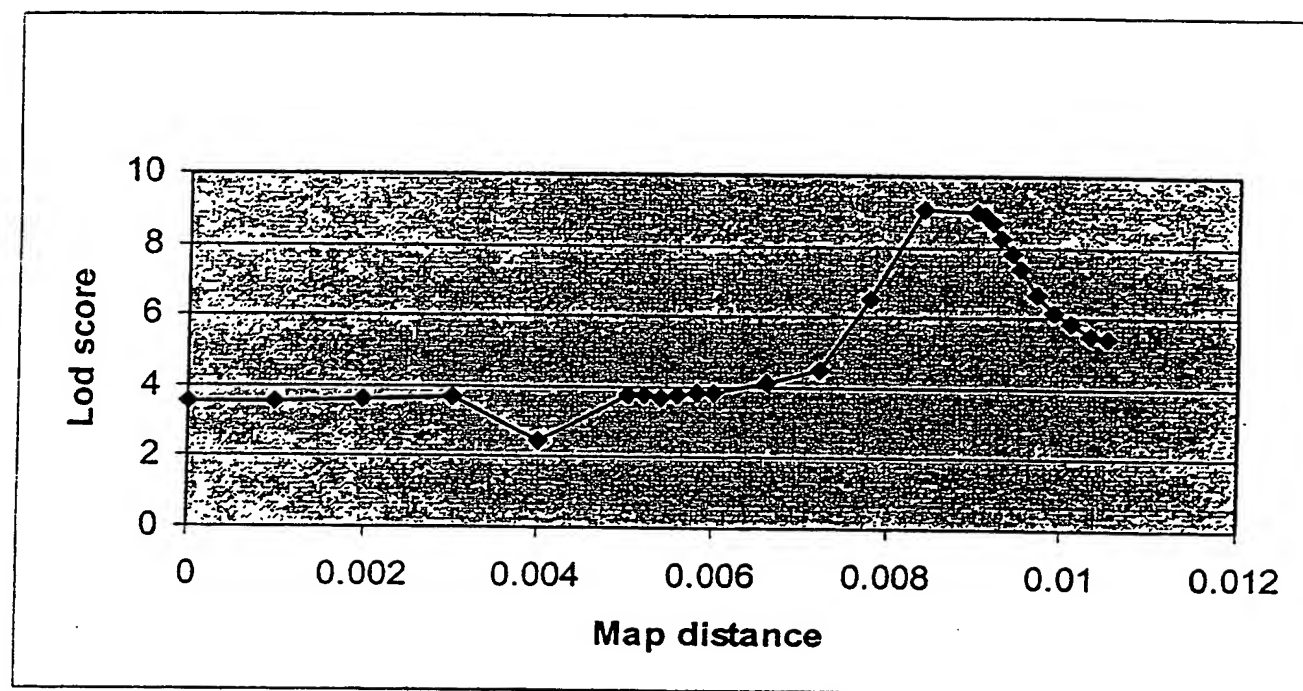


Figure 2. Alignment of pig and human *PRKAG3* cDNA sequences and deduced amino acid sequences

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Pig	TTCCTAGAGCAAGGAGAGAGCCGTTTCATGGCCATCCCGAGCTGTAACCACCAGCTCAGAAAGAAGCCATGGGGACCAGGG								
Hum	-----A-A-C-----A-C-----G-----T-G-----A-A-G-A-								
Pig	GAACAAGGCCTCTAGATGGACAAGGCAGGAGGATGTAGAGGAAGGGGGCCCTCCGGGGCCCGAGGGAAGGTCCCCAGTCCA	90	100	110	120	130	140	150	160
Hum	-GC---A---TG-----A---TCG--G-----A---A---T-A-G-----G-----								
Pig	GGCCAGTTGCTGAGTCCACCGGGCAGGAGGCCACATTCCCCAAGGCCACACCCCTTGGCCAAGCCGCTCCCTTGGCCGAG	170	180	190	200	210	220	230	240
Hum	-----AC-----T-----A-----T-----T-A---...T---G---								
Pig	GTGGACAACCCCCAACAGAGCGGGACATCCTCCCCTCTGACTGTGCAGCCTCAGCCTCCGACTCCAACACAGACCATCT	250	260	270	280	290	300	310	320
Hum	-----G---CT---A-----G-T-----TG-----A-----TG-A-G-----G-----TG--G-								
Pig	GGATCTGGGCATAGAGTTCTCAGCCTCGGCGGCGTCGGGGGATGAGCTTG...GGCTGGTGAAGAGAAGCCAGCCCCGT	330	340	350	360	370	380	390	400
Hum	---G---C---CG-----C-----A-A-A---C-G-A-TG-----A-AAG-C---C-----G---T---T---								
Pig	GCCCATCCCCAGAGGTGCTGTTACCCAGGCTGGGCTGGGATGATGAGCTGCAGAAGCCGGGGGCCAGGTCTAC	410	420	430	440	450	460	470	5' UTR
Hum	---TG-----GC---CC-CA--T---A-----C---A---G---A---C---C-----A-----								
	CDS								
Pig	ATG CAC TTC ATG CAG GAG CAC ACC TGC TAC GAT GCC ATG GCG ACC AGC TCC AAA CTG GTC	10							20
Hum	Met His Phe Met Gln Glu His Thr Cys Tyr Asp Ala Met Ala Thr Ser Ser Lys Leu Val								
	- - - - - Arg - A - - T - - - - - G - - A - - - -								
Pig	ATC TTC GAC ACC ATG CTG GAG ATC AAG AAG GCC TTC TTT GCC CTG GTG GCC AAC GGC GTC	30							40
Hum	Ile Phe Asp Thr Met Leu Glu Ile Lys Lys Ala Phe Phe Ala Leu Val Ala Asn Gly Val								
	- T - - - - - - - - - - - T - - G - -								
Pig	CGA GCG GCA CCT TTG TGG GAC AGC AAG AAG CAG AGC TTC GTG GGG ATG CTG ACC ATC ACA	50							60
Hum	Arg Ala Ala Pro Leu Trp Asp Ser Lys Lys Gln Ser Phe Val Gly Met Leu Thr Ile Thr								
	--G --A --C --- C-A --- - - - - - - - - - - - T - - - - - - - - - - - T - -								
Pig	GAC TTC ATC TTG GTG CTG CAC CGC TAT TAC AGG TCC CCC CTG GTC CAG ATC TAC GAG ATT	70							80
Hum	Asp Phe Ile Leu Val Leu His Arg Tyr Tyr Arg Ser Pro Leu Val Gln Ile Tyr Glu Ile								
	- - - - - C - - - - - T - - - - - C - - - - - - - - - - - - - - - T - - - - -								
Pig	GAA GAA CAT AAG ATT GAG ACC TGG AGG GAG ATC TAC CTT CAA GGC TGC TTC AAG CCT CTG	90							100
Hum	Glu Glu His Lys Ile Glu Thr Trp Arg Glu Ile Tyr Leu Gln Gly Cys Phe Lys Pro Leu								
	- - - C - - - - - - - - - - - - - - - - - G - - - - - - - - - - - - - - -								
Pig	GTC TCC ATC TCT CCC AAT GAC AGC CTG TTC GAA GCT GTC TAC GCC CTC ATC AAG AAC CGG	110							120
Hum	Val Ser Ile Ser Pro Asn Asp Ser Leu Phe Glu Ala Val Tyr Ala Leu Ile Lys Asn Arg								
	- - - - - T - - - - - T - - - - - T - - - - - - - - - - - A - - - - - - - - - - -								
Pig	ATC CAC CGC CTG CCG GTC CTG GAC CCT GTC TCC GGG GCT GTG CTC CAC ATC CTC ACA CAT	130							140
Hum	Ile His Arg Leu Pro Val Leu Asp Pro Val Ser Gly Ala Val Leu His Ile Leu Thr His								
	- - - T - - - - - T - - T - - T - - - - - G - - G - - A - - C AAC - - A - - - - - - - - - - -								
Pig	AAG CGG CTT CTC AAG TTC CTG CAC ATC TTT GGC ACC CTG CTG CCC CGG CCC TCC TTC CTC	150							160
Hum	Lys Arg Leu Leu Lys Phe Leu His Ile Phe Gly Thr Leu Leu Pro Arg Pro Ser Phe Leu								
	- - A - - C - - G - - - - - - - - - - - - - - - T - - T - - - - - - - - - - - - - - -								

Printed:25-01-2001

10-09-1999

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DRAW

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Pig							
Hum	ACCCCTGAGAATGAGCAATTGAGAAAACAAAACAAAAGGAACAATCCATGAACTTAGATTTTATTGGTTTCACTCAAAT							
	730	740						
Pig							
Hum	GCTGCAGTCATTGACCTG							

Figure 3. Alignment of A₁KG/SNF4 amino acid sequences from mouse, rat, human, pig, *Drosophila* and yeast

	10	20	30	40	50	60
HumG1	METVISSDSS	PAVENEHPQE	TPESNNSVYT	SFMKSHRCYD	LIPTSSKLVV	FDTSLQVKKA
RatG1	--S-.AAE-A	--P---S--	-----S----	T-----	-----	-----
MusG1	--S-.AAE--	--L---F--	-----	-----	-----	-----
HumG2						
PigG3			M H--QE-T--	AMA-----I	---M-EI--	
HumG3			M R--QE-T--	AMA-----I	---M-EI--	
Dros	RDSRGLPVAD	-FL-KVNLS	LE-DDSQIFV	K-FRF-K--	-----A--	---Q-L----
SNF4	MKPTQ	DSQ-KVSIEQ	QLAVES..IR	K-LN-KTS--	VL-V-YR-I-	L----L---S
	70	80	90	100	110	120
HumG1	FFALVTNGVR	AAPLWDSKKQ	SFVGMLTITD	FINILHRYRK	SALVQIYELE	EHKIETWREV
RatG1	-----	-----	-----	-----	-----	-----
MusG1	-----	-----	C-----	-----	-----	-----
HumG2	-----A----	-----E--T-	-SL-----	-----	-PM-----	-----L
PigG3	-----A----	-----	-----	--LV-----R	-P-----I-	-----I
HumG3	-----A----	-----	-----	--LV-----R	-P-----I-	Q-----I
Dros	-Y---Y----	-----E--	Q-----	--K--QM--	-PNASMEQ--	---LD---D-
SNF4	LNV-LQ-SIV	S-----TS	R-A-L--T--	---VIQY-FS	NPD.KFELVD	KLQLDGLKDI
	130	140	150	160	170	180
HumG1	YLQDSFKPL.	.VCISPNASL	FDAVSSLIRN	KIHRLPVIDP	ESGN....TL	YILTHKRILK
RatG1	-----	-----	-----	-----	-----	-----
MusG1	-----	-----S	-----	-----	-----	-----
HumG2	---ET-----	--N--D--	---Y--K-	-----	I-----A-	-----
PigG3	---GC-----	--S--D--	-E--YA--K-	R-----L-	V--A--V-	H-----L--
HumG3	---GC-----	--S--D--	-E--YT--K-	R-----L-	V-----V-	H-----L--
Dros	.-HNQVM--	--S-G-D--	Y--IKI--HS	R-----	AT-----V-	-----R
SNF4	ERALGVDQ-D	TAS-H-SRP-	-E-CLKMLES	RSG-I-L--Q	DEETHREIVV	SV--QY----
	190	200	210	220	230	240
HumG1	FLKLFITEFP	KPEFMSKSLE	ELQIGTYANI	AMVRTTTPVY	VALGIFVQHR	VSALPVVDEK
RatG1	-----	-----	-----	-----	-----	-----
MusG1	-----I--	-----Q	-----	-----	-----	-----
HumG2	--Q--MSDM-	--A--KQN-D	--G---H--	-FIHPD--II	K--N--ER-	I-----S
PigG3	--HI-G-LL-	R-S-LYRTIQ	D-G---FRDL	-V-LE-A-IL	T--D--DR-	-----N-T
HumG3	--HI-GSLL-	R-S-LYRTIQ	D-G---FRDL	-V-LE-A-IL	T--D--DR-	-----N-C
Dros	--F-Y-N-L-	--AY-Q--R	--K---N--	ETADE--SII	T--KK--ER-	-----L--SD
SNF4	-VA-NCR--..	.TH-LKIPIG	D-N-I-QD-M	KSCQM----I	DVIQMLT-G-	--SV-II--N
	250	260	270	280	290	300
HumG1	GRVVDIYSKF	DVINLAAEKT	YNNLDVSVTK	ALQHRSHYFE	GVLKCYLHET	LETIINRLVE
RatG1	-----	-----	-----	-----	-----	-----A-----
MusG1	-----	-----	-----	-----	-----	-----
HumG2	-K-----	-----	---IT--Q	-----Q--	--V--NKL-I	----VD-I-R
PigG3	-Q--GL--R-	---H---QQ-	--H--MN-GE	--RQ-TLCL-	---S-QP--	-GEV-D-I-R
HumG3	-Q--GL--R-	---H---QQ-	--H--M--GE	--RQ-TLCL-	---S-QP--S	-GEV-D-IAR
Dros	--L---A--	-----	--D---LR-	-NE--NEW--	--Q--N-D-S	-Y--ME-I-R
SNF4	-YLINV-EAY	--LG-IKGGI	--D-SL--GE	--MR--DD--	--YT-TKNDK	-S--MDNIRK
	310	320	330	340		
HumG1	AEVHRLVVVD	ENDVVKGIVS	LSDILQALVL	TGGEKKP*		
RatG1	-----	-H-----	-----	-----		
MusG1	-----	-H??-----	-----D--	-----		
HumG2	-----N	-A-SIV--I-	-----I-	-PAGA-QKET	ETE*	
PigG3	EQ-----L--	-TQHLL-V--	-----	SPAGIDALGA	*	
HumG3	EQ-----L--	-TQHLL-V--	-----	SPAGIDALGA	*	
Dros	-----	--RK-I--I-	-----LY--	RPSGEGV		
SNF4	-R---FF---	DVGRLV-VLT	-----KYIL-	GSN*		

Figure 4. Neighbor-joining tree constructed for mammalian, *Drosophila* and yeast AMPKG/SNF4 amino acid sequences.

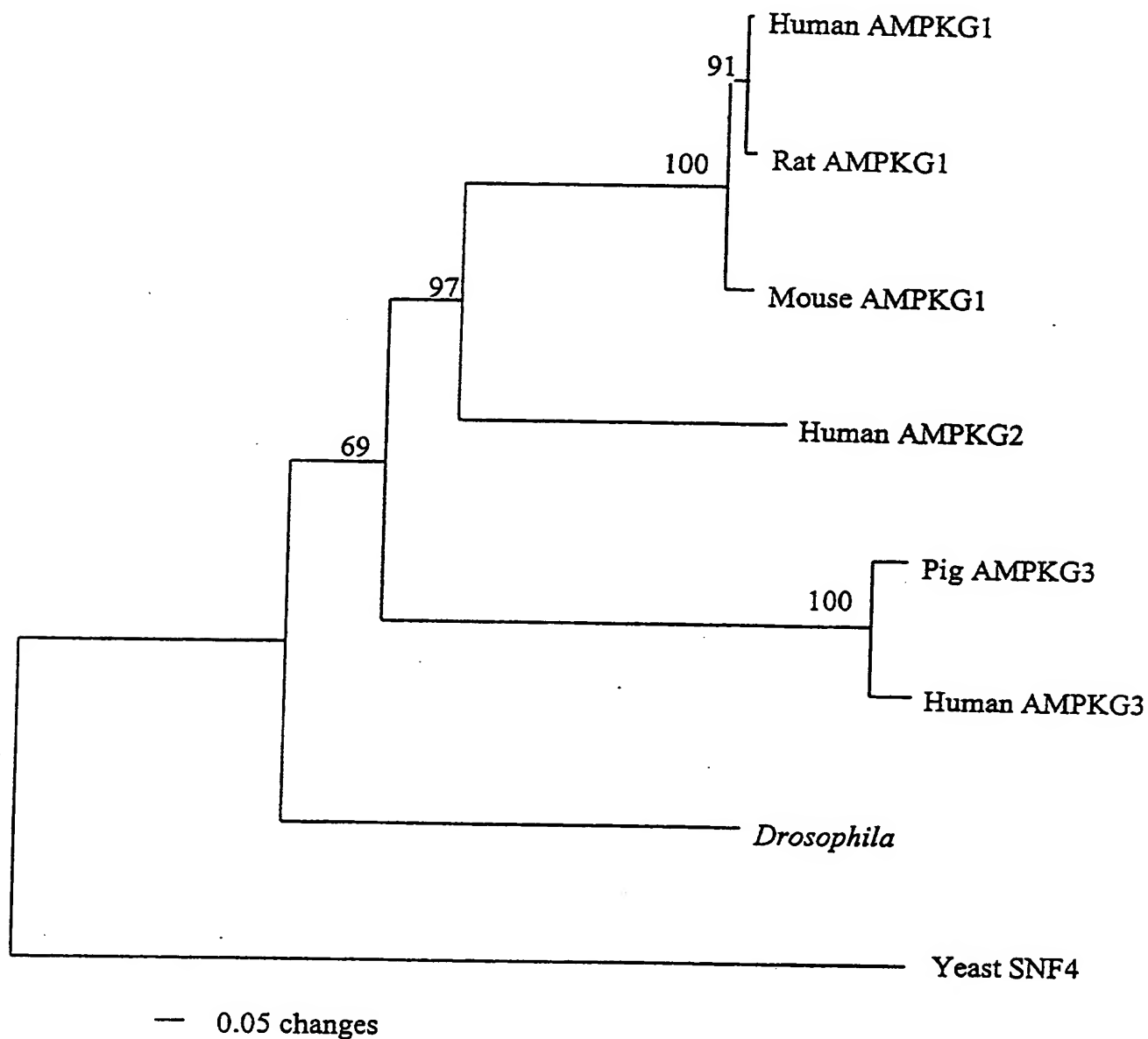


Figure 5. Northern blot analysis of human mRNA using human *PRKAG1*, human *PRKAG2* and porcine *PRKAG3* probes

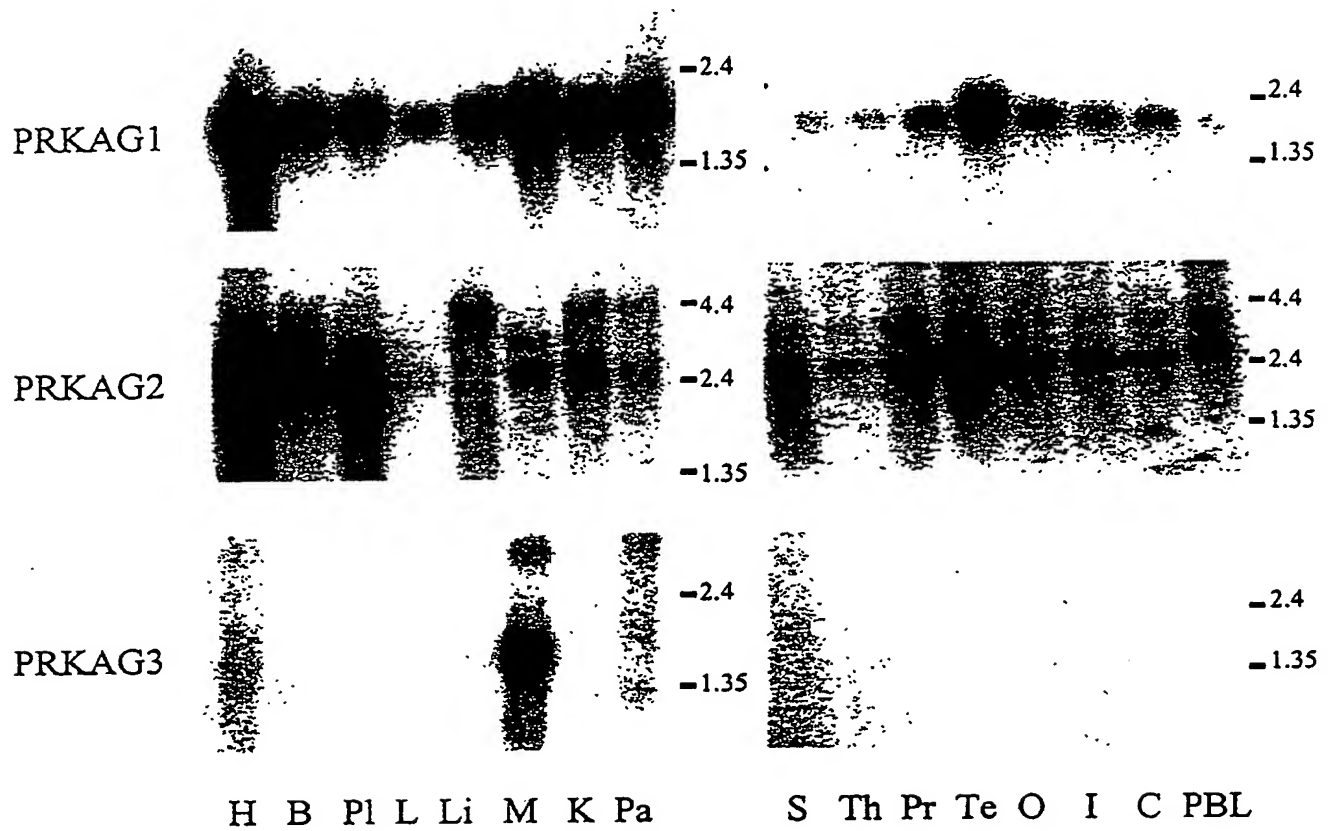
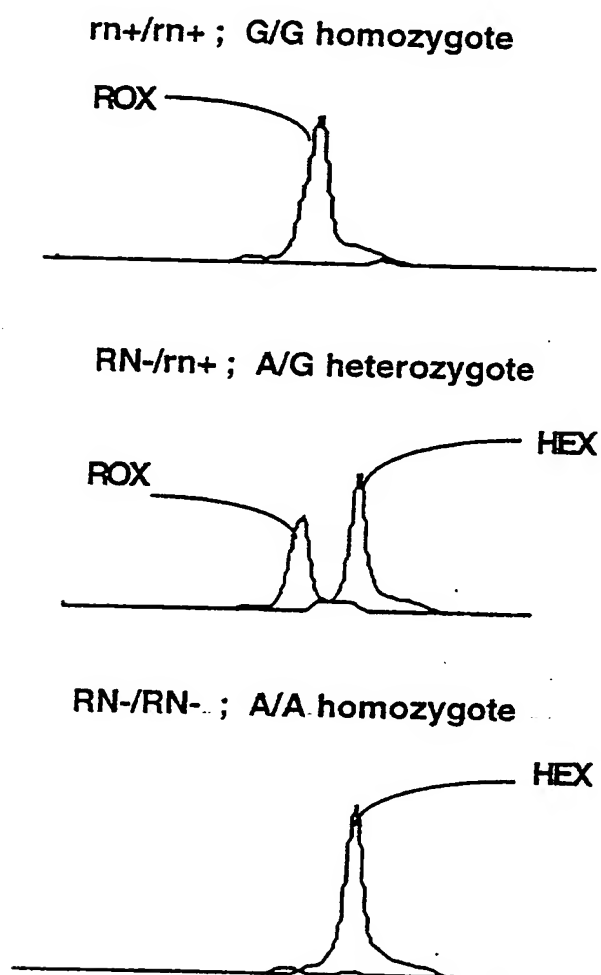


Figure 6. Detection of the missense mutation at nucleotide position 122 in *PRKAG3* associated with *RN-* allele in pigs using the OLA method.



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